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INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 6:		(11) International Publication Number: WO 95/26196
A61K 35/76, 39/12, 39/395, C07K 14/005, 16/08, C12N 1/21, 5/10, 15/33	A1	(43) International Publication Date: 5 October 1995 (05.10.95)
(21) International Application Number: PCT/US (22) International Filing Date: 29 March 1995 (20)		CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT,
(30) Priority Data: 219,262 29 March 1994 (29.03.94)	τ	Published S With international search report.
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(54) Title: CHIMERIC INFECTIOUS BURSAL DISEASED THEREON	SE VII	US cDNA CLONES, EXPRESSION PRODUCTS AND VACCINES

BASED THEREON

(57) Abstract

Chimeric cDNA for the expression of immunogenic polypeptides include the genetic epitopic determinants for a base infectious bursal disease virus strain and at least one other infectious bursal disease virus strain. The genetic epitopic determinants encode amino acids or amino acid sequences which define epitopes bound to by previously established monoclonal antibodies. The immunogens expressed by the cDNA may be employed to provide a vaccine against a plurality of IBDV strains. The epitopic determinant of IBDV lethal strains has been detected, and an immunogen for conferring immunity with respect thereto is disclosed. Similarly, a monoclonal antibody specific for IBDV lethal strains is identified, and a vaccine for passive immunization therewith is also disclosed. Immunogens exhibiting conformational epitopes, in the form of virus-like particles, are effective in the preparation of vaccines.

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Chimeric cDNA for the expression of immunogenic polypeptides include the genetic epitopic determinants for a base infectious bursal disease virus strain and at least one other infectious bursal disease virus strain. The genetic epitopic determinants encode amino acids or amino acid sequences which define epitopes bound to by previously established monoclonal antibodies. The immunogens expressed by the cDNA may be employed to provide a vaccine against a plurality of IBDV strains. The epitopic determinant of IBDV lethal strains has been detected, and an immunogen for conferring immunity with respect thereto is disclosed. Similarly, a monoclonal antibody specific for IBDV lethal strains is identified, and a vaccine for passive immunization therewith is also disclosed. Immunogens exhibiting conformational epitopes, in the form of virus-like particles, are effective in the preparation of vaccines. Data supplied from the esp@cenet database - Worldwide

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Description

Chimeric Infectious Bursal Disease Virus cDNA Clones, Expression Products and Vaccines Based Thereon

Technical Field:

The present invention provides chimeric IBDV immunogens which actively protect against virulent and lethal challenge by Classic and variant IBDV strains, and methods for obtaining vaccines containing these chimeric immunogens and vaccines.

Background Art

Infectious bursal disease virus (IBDV) is responsible for a highly contagious immunosuppressive disease in young chickens which causes significant losses to the poultry industry worldwide (reviewed in <u>Kibenge</u> (1988) "J. Gen. Virol.", 69:1757-1775). Infection of susceptible chickens with virulent IBDV strains can lead to a highly contagious immunosuppressive condition known as infectious bursal disease (IBD). Damage caused to the lymphoid follicles of the bursa of *Fabricius* and spleen can exacerbate infections caused by other agents and reduce a chicken's ability to respond to vaccination as well (<u>Cosgrove</u> (1962) "Avian Dis.",

There are two serotypes of IBDV (McFerran et al (1980) "Avian Path." 9:395-404). Serotype 1 viruses are pathogenic to chickens and differ markedly in their virulence (Winterfield et al (1978) "Avian Dis." 5:253-260), whereas serotype 2 viruses, isolated from turkeys, are avirulent for chickens (Ismail et al (1988) "Avian Dis.", 32:757-759; Kibenge (1991) "Virology" 184:437-440).

IBDV is a member of the *Birnaviridae* family and its genome consists of two segments of double-stranded RNA (<u>Dobos et al</u> (1979) "J. Virol.", 32:593-605). The smaller segment B (~2800bp) encodes VP1, the dsRNA polymerase. The larger genomic segment A (~3000bp) encodes a 110 kDa precursor polyprotein in a single open reading frame (ORF) that is processed into mature VP2, VP3 and VP4 (<u>Azad et al</u> (1985)

"Virology" 143:35-44). From a small ORF partly overlapping with the polyprotein ORF, segment A can also encode VP5, a 17 Kda protein of unknown function (<u>Kibenge et al</u> (1991) "J. Gen. Virol.", 71:569-577).

While VP2 and VP3 are the major structural proteins of the virion, VP2 is the major host-protective immunogen and causes induction of neutralizing antibodies (Becht et al (1988) "J. Gen. Virol." 69:631-640; Fahey et al (1989) "J. Gen. Virol.", 70:1473-1481). VP3 is considered to be a group-specific antigen because it is recognized by monoclonal antibodies (Mabs) directed against VP3 from strains of both serotype 1 and 2 (Becht et al (1988) "J. Gen. Virol.", 69:631-640). VP4 is a virus-coded protease and is involved in the processing of the precursor protein (Jagadish et al (1988) "J. Virol.", 62: 1084-1087).

In the past, control of IBDV infection in young chickens has been achieved by live vaccination with avirulent strains, or principally by the transfer of maternal antibody induced by the administration of live and killed IBDV vaccines to breeder hens. Unfortunately, in recent years, virulent variant strains of IBDV have been isolated from vaccinated flocks in the United States (Snyder et al (1988b) "Avian Dis.", 32:535-539; Van der Marel et al (1990) "Dtsch. Tierarztl. Wschr.", 97:81-83). The use of a select panel of Mabs, raised against various strains of IBDV, has led to the identification of naturally occurring GLS, DS326, RS593 and Delaware variant viruses in the United States. Substantial economic losses have been sustained due to the emergence of these antigenic variants (Delaware and GLS) in the field (Snyder et al (1992) "Arch. Virol.", 127:89-101), copending U.S. Application Serial No. 08/216,841, filed March 24, 1994, Attorney Docket No. 2747-053-27, Snyder, copending herewith). These variant strains are antigenically different from the Classic strains of IBDV most typically isolated before 1985, and lack epitope(s) defined by neutralizing monoclonal antibodies

(Mabs) B69 and R63 (Snyder et al (1988a) "Avian Dis.", 32:527-534; Snyder et al (1998b) "Avian Dis.", 32:535-539; Snyder et al (1992) "Arch. Virol.", 127:89-101). Since the appearance of these variant strains in the field, many commercially available live and killed vaccines for IBDV have been reformulated in an attempt to better match the greater antigenic spectrum of viruses recognized to be circulating in the field.

been made, and the genome of IBDV has been cloned (Azad et al (1985) "Virology", 143:35-44). The VP2 gene of IBDV has been cloned and expressed in yeast (Macreadie et al (1990) "Vaccine", 8:549-552), as well as in a recombinant fowlpox virus (Bayliss et al (1991) "Arch. Virol.", 120:193-205). When chickens were immunized with the VP2 antigen expressed from yeast, antisera afforded passive protection in chickens against IBDV infection. When used in active immunization studies, the fowlpox virus-vectored VP2 antigen afforded protection against mortality, but not against damage to the bursa of Fabricius.

Recently, the synthesis of VP2, VP3 and VP4 structural proteins of the variant GLS IBDV strain in a baculovirus expression system has been described (Vakharia et al (1993) "J. Gen. Virol.", 74:1201-1206). In an initial two dose active immunity study in SPF chickens, baculovirus expressed GLS proteins were able to confer 79% protection against virulent GLS challenge (Vakharia et al (1993) "J. Gen. Virol.", 74:1201-1206). In a subsequent extended study of active cross-immunity, by increasing the antigenic mass of the baculovirus expressed GLS protein, complete protection against the variant GLS and E/Del strains was obtained with a single dose, but only partial protection was afforded against the Classic STC strain unless two doses were administered.

In recent years, the complete, nucleotide sequences of the large segment A of five serotype 1 IBDV strains; 002-73

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(<u>Hudson et al</u> (1986) "Nucleic Acids Res." 14:001-5012), Cu-1, PBG98, 52/70 (<u>Bayliss et al</u> (1990) "J. Gen. Virol.", 71:1303-1312), STC (<u>Kibenge</u> (1990) "J. Gen. Virol.", 71:569-577), and serotype 2 OH strain (<u>Kibenge</u> (1991) "Virology", 184:437-440) have been determined. In addition, the VP2 gene of virulent Japanese IBDV strains (<u>Lin et al</u> (1993) "Avian Dis.", 37:315-323) and Delaware variants A and E (<u>Lana et al</u> (1992) "Virus Genes" 6:247-259; <u>Heine et al</u> (1991) "J. Gen. Virol.", 22:1835-1843) has been sequenced. However, noone has completely cloned and characterized the entire long segment of any United States IBDV variant.

Disclosure of the Invention

Inventors have now identified the region of the IBDV genome which is responsible for antigenic variation. A DNA sequence containing the central variable region of VP2 protein, as well as a plasmid incorporating the same, have been constructed. This DNA sequence can be manipulated to generate desired virus neutralizing epitopes or immunogenic polypeptides of any IBDV strain. In turn, these immunogenic segments can be incorporated into new recombinant IBDV vaccines.

Brief Description of the Drawings

Figure 1 illustrates the construction of various chimeric plasmids encoding IBDV-specific polyproteins. A map of the IBDV genome with its coding regions is shown at the top of the Figure. Selected restriction sites are incorporated in the Figure: B, BamHI; E, BstEII; N, NdeI; R, NarI; S, SpeI. Dashes indicated the substitution of the D78 sequence (NdeI-NarI fragment) into the GLS sequence to restore the B69 epitope region. Solid line and dotted line indicate the substitution of the E/Del-22 and DS326 sequences, respectively, into the GLS sequence to restore the B63 epitope region or to delete the 179 epitope region, respectively.

Figure 2 is electron micrographs of IBDV virus-like particles (|---|) = 100nm). A. Actual empty particles (without RNA) from purified virus. B. Virus-like particles (empty capsids) derived from a recombinant baculovirus expressing the large genome segment of IBDV in insect cells

Figure 3 is a comparison of the deduced amino acid sequences of the structural proteins (VP2, VP3 and VP4) of ten IBDV strains. Dashes (-) indicate amino acid identity and crosses (x) denote a region where the sequence was not determined. Filled bar (1) indicates a gap in the sequence and vertical arrowheads (1) mark the possible cleavage sites of VP2/VP4 and VP4/VP3. The two hydrophilic peaks in the variable region are overlined.

Figure 4 is a phylogenetic tree for the IBDV structural proteins using the PAUP (phylogenetic analysis using parsimony) version 3.0 program (Illinois Natural History Survey, Champaign, Illinois).

Figure 5 reflects the DNA and amino acid sequence for the GLS virus structural protein fragment VP2/VP4/VP3. A vertical line indicates the start/stop points for the VP2, VP4 and VP3 regions.

Figure 6 reflects the DNA and amino acid sequence for the E/Del 22 virus structural protein fragment VP2/VP4/VP3.

Figure 7 is a table of the amino acid identities for key locations (epitopic determinants) of eight different IBDV.

Definitions:

IBD - infectious bursal disease as described above.

<u>IBDV</u> - infectious bursal disease virus, a virus capable of, at a minimum, inducing lesions in the bursa of *Fabricius* in infected poultry.

<u>Epitopic Determinants</u> - amino acids or amino acid sequences which correspond to epitopes recognized by one or more monoclonal antibodies. Presence of the amino acid or amino acid sequence at the proper ORF location causes the

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polypeptide to exhibit the corresponding epitope. An epitopic determinant is identified by amino acid(s) identity and sequence location.

<u>Genetic Epitopic Determinants</u> - nucleotide sequences of the ORF which encode epitopic determinants.

Conformational Epitopes - epitopes induced, in part or in whole, by the quaternary (three-dimensional) structure of an IBDV polypeptide. Conformational epitopes may strengthen binding between an IBDV and a monoclonal antibody, or induce binding whereas the same sequence, lacking the conformational epitope, would not induce binding between the antibody and the IBDV polypeptide at all.

<u>Virus-Like Particles</u> - three-dimensional particles of natural or recombinant amino acid sequences mimicking the three-dimensional structure of IBDV (encoded by the large genome segment A) but lacking viral RNA. Virus-like particles exhibit conformational epitopes exhibited by native viruses of similar sequence. Virus-like particles are created by the proper expression of DNA encoding VP2, VP4, VP3 structural proteins in a proper ORF.

Epitopic Determinant Region - Limited region of amino acid sequence of VP2 of IBDV that is replete with epitopic determinants, variation among amino acids of this limited region giving rise to a high number of epitopes recognized by different monoclonal antibodies.

Best Mode for Carrying Out the Invention

Recombinant, immunogenic polypeptides exhibiting the epitopes of two or more native IBDV, as well as recombinant virus-like particles exhibiting the epitopes of two or more native IBDV and conformational epitopes are effective immunogens for vaccines which can be used to confer protection against a wide variety of IBDV challenge in inoculated poultry. The recombinant polypeptides and virus-like particles are obtained by the expression of chimeric DNA

prepared by the insertion, in the VP2 region of a base IBDV, of epitopic determinants for at least a second IBDV. most easily done by substitution of the genetic epitopic determinants for the amino acids identities and locations reflected in Figure 7. Thus, where the epitopic determinant of the second IBDV differs from that of the base IBDV, the genetic epitopic determinant for the differing second IBDV is inserted in place of the genetic epitopic determinant at that location of the base IBDV. An example, combining epitopic determinants from the D78, E/Del 22 and DS326 IBDV into the base GLS IBDV is set forth in Figure 1. Thus, one DNA sequence can be prepared with genetic epitopic determinants for a plurality of native IBDV. These recombinant plasmids can be inserted into a variety of packaging/expression vector, including baculovirus, fowlpox virus, Herpes virus of turkeys, adenovirus and similar transfection vectors. The vectors can be used to infect conventional expression cells, such as SF9 cells, chicken embryo fibroblast cell lines, chicken embryo kidney cells, vero cells and similar expression vehicles. Methods of transfection, and methods of expression, as well as plasmid insertion into transfection vehicles, are well known and do not constitute an aspect of the invention, per se.

The expression of the chimeric cDNA of the invention generate immunogenic polypeptides which reflect epitopes of a plurality of native IBDV, and the expression of a recombinant VP2, VP4, VP3 cDNA segment, with the VP2 region again comprising genetic epitopic determinants for at least two native IBDV give rise to immunogenic virus-like particles.

The immunogenic polypeptides and virus-like particles can be harvested using conventional techniques (<u>Dobos et al</u>, "J. Virol.", 32:593-605 (1979)). The polypeptides and virus-like particles can be used to prepare vaccines which will confer protection on inoculated poultry, in particular, chickens, and in a preferred embodiment, broiler chickens, protection against challenge from each IBDV bearing an epitope reflected

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in the plurality of epitopic determinants present in the inoculum. Thus, a single immunogen gives rise to immunity against a variety of IBDV, each IBDV whose genetic epitopic determinant has been incorporated in the chimeric cDNA.

The administration of the vaccines can be effectively done according to well-established procedures. In U.S. Patent 5,064,646, which is incorporated herein by reference, methods are described for the effective inoculation of chicks based on the then novel isolation of GLS IBDV. Similar administration and dosage regimens can be employed herein. Since the polypeptides and virus-like particles lack viral RNA, they are avirulent. The vaccines may therefor be prepared by simple incorporation of the immunogenic polypeptides and virus-like particles in a pharmaceutical carrier, typically a suspension or mixture. Appropriate dosage values are best determined through routine trial and error techniques, given the different antibody titers induced and/or the quantity of different epitopes present which will induce complete crossimmunity to virulent challenge. In general, pharmacologically acceptable carriers such as a phosphate buffered saline, cell culture medium, Marek's virus vaccine diluent oil adjuvants and other adjuvants, etc., can be used. Administration is preferably done to hens entering egg laying periods which provides induction of antibody which is passively transferred through the egg to the chick to prevent early invention by virulent field strength IBDV. Conversely, the recombinant vaccine may be delivered in a replicating vector at any time in a chicken's life span, preferably at one day of age. Experience has demonstrated that, generally, that the level of protection may be improved by a second inoculation.

This invention may be further understood by reference to the specific examples set forth below.

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Examples:

Background Methodology

To determine the molecular basis of antigenic variation in IBDV, the genomic segment A of four IBDV strains: GLS, DS326, Delaware variant E (E/Del) and D78 was cloned and characterized by sequencing. By comparing the deduced amino acid sequences of these strains with other serotype 1 and 2 sequences published previously, the putative amino acid residues involved in the binding with various neutralizing Mabs were identified, and the phylogenetic relationship of IBDV structural proteins was examined.

GLS, DS326 and STC strains of IBDV were propagated in the bursa of specific-pathogen-free chickens (SPAFAS, Inc., Norwich, CT, USA). Tissue culture adapted E/Del-22, D78 and OH (serotype 2) strains of IBDV were propagated in primary chicken embryo fibroblast cells derived from 10-day-old embryonated eggs (SPAFAS, Inc.) and purified as described (Snyder et al (1988a) "Avian Dis.", 32:527-534). against various strains of IBDV were produced and characterized using protocols previously outlined (Snyder et al (1988a) "Avian Dis." 32:527-534; Snyder et al (1988b) "Avian Dis.", 32:535-539). Mabs B69 and R63 were prepared against D78 strain, whereas Mabs 8, 10, 57 and 179 were prepared against GLS strain. In addition, a new Mab 67 was prepared which was neutralizing and specific for the E/Del strain. Identification of IBDV antigens by modified antigen capture ELISA (AC-ELISA) was carried out as described (Snyder et al (1992) "Arch. Virol.", 127:89-101).

Various strains of IBDV were characterized by their reactivities with a panel of neutralizing Nabs, as shown in Table 1.

TABLE 1

Antigenic characterization of various IBDV strains by their reactivities with a panel of neutralizing MAbs

				Reactivities with MAbs	ties w	th Mab	. 89		
Virus Strains	Classification	B69	R63	179	8	10	57	67	
D78	Classic	+	+	+	+	+	1	ı	
PBC98	Classic	i,	+ -	+	+	+	ı	I,	
STC	Classic	+	+	+	+	+	i	ı	
52/70	Classic	+	+	+	+	1	ı	1	
OH (serotype 2)	Classic	+	+	+	+	ı	1	1	
E/Del	Variant	ı	+	+	+	ı	ı	+	
STS	Variant	1	i	+	+	+	+	i	
DS326	Variant	ı	ı	1	+	+	+	1	

All standard serotype 1 viruses reacted with Mabs B69, R63, 179 and 8, except PBG98 (a British vaccine strain, Intervet, U. K.) which did not react with Mab B69. In contrast, all the U.S. variant viruses lack the virus-neutralizing B69 epitope. In addition, GLS and DS326 variants lack an R63 epitope but share a common epitope defined by the Mab 57. Thus, on the basis of the reactivities with various Mabs, these viruses were antigenically grouped as classic, GLS, DS326 and E/Del variants.

Complementary DNA clones, containing the entire coding region of the large RNA segment of various IBDV strains, were prepared using standard cloning procedures and methods previously described (<u>Vakharia et al</u> (1992) "Avian Dis.", 36:736-742; <u>Vakharia et al</u> (1993) "J. Gen. Virol.", 74:1201-1206). The complete nucleotide sequence of these cDNA clones was determined by the dideoxy method using a Sequenase DNA sequencing kit (U.S. Biochem. Corp., Columbus, OH). DNA sequences and deduced amino acid sequences were analyzed by a PC/GENE software package (Intelligenetics, Inc.). These are reflected in Figures 5 and 6. The nucleotide sequence data of the GLS strain has been deposited with GenBank Data Libraries and has been assigned an accession number M97346.

Comparisons of the nucleotide sequence of GLS strain (3230 bp long) with eight serotype 1 and one serotype 2 IBDV strains exhibit ≥ 92% and ≥ 82% sequence homology, respectively; indicating that these viruses are closely related. It is interesting to find that there are only six to nine base substitutions between D78, PBG98, and Cul strains which corresponds to a difference of about 0.2% to 0.3% (results not shown). Figure 3 and Table 2 show a comparison of the deduced amino acid sequences and percent homology of the large ORF of segment A of the ten IBDV strains, including four IBDV strains used in this study.

TABLE 2

9	ОН										
Percent amino acid sequence homology of large Okk of Begment A Of ten laby strains	002-73										•
מנ רפוו	STC									97.4	
gment A	PBG98 52/70								98.3	97.3	6
KF OI BE	PBG98							98.3	98.3	97.6	0
large u	Cu-1						99.5	98.5	98.5	7.76	0
ogy or	D78					9.66	99.5	98.4	98.4	9.76	
ce nomol	E/Del				97.9	0.86	97.9	97.9	97.5	7.96	
d seguen	DS326			98.3	98.1	98.2	98.1	98.1	98.0	97.1	
umino aci	GLS		7.86	98.4	98.5	98.6	98.5	98.1	7.76	97.0	
Percent (Strain	GLS	DS326	E/Del	D78	Cu-1	PBG98	52/70	STC	002-73	

These comparisons show that the proteins are highly conserved. The degree of difference in the amino acid sequence ranges from 0.4% for the D78 versus Cu-1 comparison and 10.3% for the serotype 1 (E/Del) versus serotype 2 (OH) comparison (Table 2).

In Figure 3, alignments of the deduced amino acid sequences of the large ORF (1012 residues) of ten IBDV strains (including four used in this study) show that most of the amino acid changes occur in the central variable region between residues 213 and 332 of VP2 protein, as shown earlier by <u>Bayliss</u> et al (1990) "J. Gen. Virol. M, 71:1303-1312. is interesting to note that all the U.S. variants (GLS, DS326 and E/Del) differ from the other strains in the two hydrophilic regions which are overlined in Figure 3 (residues 212 to 223 and residues 314 to 324). These two hydrophilic regions have been shown to be important in the binding of neutralizing Mabs and hence may be involved in the formation of a virus-neutralizing epitope (Heine et al (1991) "J. Gen. Virol.", 22:1835-1843). Recently, we demonstrated that the conformation dependent Mabs B69, R63, 8, 179, 10, and 57 (see Table 2) immunoprecipitate VP2 protein (Snyder et al (1992) "Arch. Virol.", 127:89-101). In addition, E/Del specific Mab 67 also binds to VP2 protein. Therefore, to identify the amino acids involved in the formation of virus-neutralizing epitopes, and hence the antigenic variation, we compared the amino acid sequences of VP2 protein of classic and variant viruses.

Comparison of the D78 sequence with the PBG98 sequence shows only four amino acid substitutions at positions 76, 249, 280 and 326. However, STC and 52/70 strains also differ from the D78 sequence at positions 76, 280 and 326 but these viruses do bind to Mab B69. This implies that Gln at position 249 (Gln249) may be involved in the binding with Mab B69. It should be noted that all U.S. variant viruses have a Gln-Lys substitution at this position and hence escape the binding

with neutralizing Mab B69. Similarly, comparison of the GLS sequence with the DS326 sequence in the variable region shows six amino acid substitutions at positions 222, 253, 269, 274, 311 and 320. However, other strains of IBDV that do bind to Mab 179 have amino acid substitutions at positions 222, 253, 269 and 274 that are conservative in nature. Therefore, this suggests that Glu311 and Gln320 may be involved in the binding with Mab 179. Again, comparison of GLS and DS326 sequences with all other IBDV sequences shows a unique Ala-Glu substitution at position 321, suggesting the contribution of this residue in the binding with Mab 57. Since Mab 57 does not compete with Mab R63, it is conceivable that Ala321 may contribute to the binding with Mab R63. Similarly, comparison of E/Del sequence with other sequences shows five unique substitutions at positions 213, 286, 309, 318 and 323. However, comparison of this E/Del sequence (from tissue culture derived virus) with previously published VP2 A/Del and E/Del sequences (bursa derived virus) suggests the involvement of Ile286, Asp318 and Glu323 in the binding with Mab 67 since residues at positions 213 and 309 are not substituted in A/Del and E/Del sequences, respectively (Heine et al (1991) "J. Gen. Virol.", 22:1835-1843; <u>Lana et al</u> (1992) "Virus Genes", 6:247-259; <u>Vakharia et al</u> (1992) "Avian Dis.", 36:736-742).

Comparisons of the amino acid sequence also show a striking difference between serotype 1 and serotype 2 sequences. In serotype 2 OH strain, there is an insertion of an amino acid residue at position 249 (serine) and a deletion of a residue at position 680. Previously, it has been shown that serotype 2 viruses are naturally avirulent and do not cause any pathological lesions in chickens (Ismail et al (1988) "Avian Dis.", 32:757-759). Thus, these subtle changes in the structural proteins of serotype 2 OH strain may play an important role in the pathogenicity of the virus. Moreover, it has been hypothesized that an amino acid sequence motif, S-W-S-A-S-G-S, (residues 326 to 332) is conserved only in

virulent strains and could be involved in virulence (Heine et al (1991) "J. Gen. Virol.", 22:1835-1843). This sequence motif was also conserved in various pathogenic strains of IBDV isolated in Japan (Lin et al (1993) "Avian Dis.", 37:315-323). comparison of the amino acid sequences in this heptapeptide region reveals that nonpathogenic serotype 2 OH strain has three substitutions, whereas mildly pathogenic strains of serotype 1 (D78, Cu-1, PBG98 and 002-73) have one or two substitutions in this region. Moreover, comparison of the hydrophilicity plots of the variable region (amino acids 213 to 332) of variant serotype 1 strains and serotype 2 OH strain indicates a drastic reduction in the second hydrophilic peak region (amino acid residues 314 to 324) for serotype 2 (results not shown). Since most of the amino acid residues causing antigenic variation reside in this region, these residues may play an important role in the formation of virusneutralizing epitopes, as well as serotype specificity.

To evaluate the antigenic relatedness of structural proteins of various IBDV strains, a phylogenetic tree was constructed, based on the large ORF sequences of ten IBDV strains, including the U.S. variant strains examined in this study (Figure 4). Three distinguishable lineages were formed. The first one, which is most distant from the others, is serotype 2 OH strain, and the second one is the geographically distant Australian serotype 1 strain (002-73). The third lineage consists of four distinct groups. The first and second group include highly pathogenic strains, namely, standard challenge (STC) strain from U. S. and the British field strain (52/70). The third group comprises all the European strains: the vaccine strains D78 (Holland), PBG98 (U.K.), and mildly pathogenic strain Cu-1 (Germany). fourth group consists of the U.S. variant strains in which E/Del forms a different subgroup. The groups formed by the phylogenetic analysis correlate very well with the Mabs reactivity patterns (see Table 1). As shown in Figure 4, all

36:736-742). To insert the chimeric IBDV structural genes in the Baculovirus genome, plasmid pB69GLS was completely digested with BstEII enzyme and partially with the BamHI enzyme, combined with the NheI-BstEII fragment (derived from plasmid pGLSBacI, see Vakharia et al (1993) "J. Gen. Virol.", 74:1201-1206) and then ligated to the NheI-BamHI cut transfer vector pBlueBacII (Invitrogen Corp., San Diego, CA). Finally, recombinant baculovirus I-7 was obtained using previously described procedures (Vakharia et al (1993) "J. Gen. Virol.", 74:1201-1206). See Table 3.

Preparation of an inoculum for immunization

Spodoptera frugiperda SF9 cells, infected at a multiplicity of 5 PFU per cell with the I-7 recombinant baculovirus, were propagated as suspension cultures in one liter flasks containing Hink's TNM-FH medium (JHR Biosciences, Lenxa, KS) supplemented with 10% fetal calf serum at 28°C for 3 to 4 days. The infected cells were recovered by low speed centrifugation, washed two times with PBS, and resuspended in a minimum volume of PBS. The cell slurry was sonicated on and ice bath three times for 1 min, with 2 min intervals and clarified by low speed centrifugation. An aliquot of each cell lysate was tested with anti-IBDV Mabs by AC-ELISA to estimate the antigenic mass present (Snyder et al (1998b) "Avian Dis.", 32:535-539). Preparations having the highest antigenic mass were pooled and comparatively titrated in AC-ELISA against the V-IBDV-7-1 recombinant baculovirus IBDV vaccine used in a previous study (Vakharia et al (1993) "J. Gen. Virol.", 74:1201-1206). The antigenic mass of the I-7 recombinant preparation, as determined by AC-ELISA with group specific neutralizing Mab 8, was adjusted by dilution to be the same as the V-IBDV-7-1 vaccine, and then it was emulsified with an equal volume of Freund's incomplete adjuvant and used for inoculation.

the U.S. variant viruses which lack the B69 epitope form a distinct group, whereas all the classic viruses containing a B69 epitope form another group (except PBG98). In addition, closely related GLS and DS326 strains containing a common Mab 57 epitope and lacking an R63 epitope could be separated from the other variant E/Del strain.

Based on this information, a recombinant vaccine was constructed as follows:

Construction of recombinant baculovirus clones containing chimeric IBDV genes

A recombinant baculovirus which expresses a chimeric VP2, VP3 and VP4 structural proteins of the GLS strain was constructed and assessed. The recombinant baculovirus expressed a chimeric VP2 protein incorporating all Mab defined GLS neutralization sites, as well as one neutralization site (B69) which is specific for Classic strains of IBDV in the form of a VP2-VP4-VP3 segment.

Complementary DNA clones, containing the entire coding region of the large RNA segment of the GLS and D78 IBDV strains, were prepared using standard cloning procedures and methods previously described (Vakharia et al (1992) "Avian Dis.", 36:736-742; Vakharia et al (1993) "J. Gen. Virol.", 74:1201-1206). To insert the gene sequence encoding the B69 epitope of the D78 IBDV strain, plasmid pB69GLS was constructed as follows (see Figure 1). Full-length cDNA clones of D78 and GLS (plasmids pD78 and pGLS-5) were digested with NdeI-NarI and NarI-SpeI enzymes to release a NdeI-NarI (0.26 kb) and a NarI-SpeI (0.28 kb) fragments, respectively. These two fragments were then ligated into the NdeI-SpeI cut plasmid pGLS-5 to obtain a chimeric plasmid pB69GLS. As a result of this insertion, three amino acids were substituted in the GLS VP2 protein. These substitutions were at positions 222 (Thr-Pro), 249 (Lys-Gln) and 254 (Ser-Gly) in the variable region of the VP2 protein (Vakharia et al (1992) "Avian Dis.",

Viruses

The challenge viruses: Classic strains IM and STC, and variant strains E/Del and GLS-5 were obtained from previously acknowledged sources (Snyder et al (1988a) "Avian Dis.", 32:527-534; Snyder et al (1992) "Arch. Virol.", 127:89-101). After intraocular instillation, challenge viruses were titrated in the bursae of specific-pathogen-free (SPF) chickens (SPAFAS, Inc., Storrs, Conn.). For strains STC, E/Del and GLS-5, a 100 chick infective fifty percent dose (100 CID₅₀) was determined based on bursa to body weight measurements. One hundred lethal doses (100 LD) of the IM strain were calculated based on mortality at 8 days postinoculation (PI).

Chicken inoculations and IBDV challenge

White leghorn SPF chickens were hatched and reared in HEPA filtered isolation units (Monair Anderson, Peachtree City, GA). Eight-week old chickens were prebled, individually wing banded, divided among 10 groups of 15 chicks each and treated as follows. Chickens of groups I-V received no inoculations and served as either negative or positive challenge controls. Chickens of group V-X were inoculated intramuscularly with 0.5 ml of the 1-7 inoculum prepared above from recombinant Baculovirus infected cell lysates. At 3 weeks PI, all chickens were bled and chickens of groups II-IX were challenged with the appropriate IBDV challenge strain by ocular instillation. Four days post-challenge, 5 chickens from each group were humanely sacrificed and their cloacal bursa were removed. Each bursa was processed and subsequently evaluated for the presence of IBDV antigen by AC-ELISA as described (Snyder et al (1998b) "Avian Dis.", 32:535-539). addition, chickens in the IM challenged groups were scored as dead, and humanely sacrificed when they became obviously moribund due to IM challenge. Eight days post-infection, the remaining chickens in all groups were sacrificed and weighed.

The bursa of Fabricius from each chicken was carefully excised and also weighed. Bursa weight to body weight ratio was calculated for each chicken as described by Lucio and Hitchner (Lucio et al (1979) "Avian Dis.", 23:466-478). Any value for individually challenged chickens falling plus or minus two standard deviation units from the mean of the corresponding control group was scored as a positive indicator of IBDV infection. Opened bursae were fixed by immersion in 10% neutral buffered formalin. Transverse portions of bursae were processed through graded alcohols and xylene, embedded into paraffin, sectioned, stained with hematoxylin-eosin, and examined with a light microscope. Protection against challenge was defined as the absence of any IBDV-induced lesions in the bursa of Fabricius.

Serological evaluation

The Classic D78 strain, as well as the cell culture adapted variant GLS strain of IBDV were grown in primary chicken embryo fibroblast cells and used in virus neutralization (VN) tests to test sera from the vaccine trial essentially as described (Snyder et al (1988a) "Avian Dis.", 32:527-534). Serum from the trials was also tested for the presence of anti-IBDV antibody using a commercially available IBDV antibody ELISA kit (Kirkegaard and Perry, Gaithersburg, MD).

Evaluation of vaccines and challenge viruses

The antigenic content of the I-7 GLS chimeric IBDV vaccine was assessed in AC-ELISA with a panel of VP2 and VP3 specific Mabs. The relative antigenic mass of each epitope expressed in the I-7 vaccine was compared to previously tested lots of Baculovirus expressed unmodified GLS subunit vaccines (Vakharia et al (1993) "J. Gen. Virol.", 74:1201-1206). The status of each Mab defined epitope on the I-7 chimeric vaccine was also compared to the status of those Mab defined epitopes

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occurring on wild type IBDV challenge viruses used to evaluate the efficacy of the I-7 vaccine. Table 3 shows that antigenic mass levels at the 8, 57, and B29 epitopes for the current I-7 chimeric vaccine were similar to a recently tested unmodified V-IBDV-7-1 GLS subunit vaccine, but approximately 4-fold higher than the original unmodified V-IBDV-7 vaccine.

TABLE 3

Comparative levels of IBDV, VP2, and VP3 monoclonal antibody (Mab) defined epitopes of recombinant baculovirus expressing IBDV proteins and status of Mab defined epitopes on challenge viruses used.

		elative]	evel of !	Relative level of Mab epitope	G.A.	Challenge		Statu	Status of Mab epitope ^B	pitope	
Vaccine	8	57	B69	67	B29	Virus	.8 _C	57 ^c	в69°	2L9	B29 ^D
V-TBD-78	1	1	0	0	1	GLS	+	+	1	1	+
V-TBD-7-18	F	9	٥	0	7	SIC	+	1	+	1	+
T-7F	, ,	8	6	0	2	IM	+	ı	+	1	+
	,					E/Del	+	1	•	+	+

The relative level of each Mab epitope was determined by AC-ELISA, and the level of each Mab epitope was set to 1 for the V-IBD-7 vaccine previously used (15). Maximum level is 9. Each 1.0 increment represents approximately twice the amount of the epitope present in the original V-IBD-7 vaccine. The V-IBD-7-1 vaccine was also previously reported (16).

The status of Mab epitopes was determined by AC-ELISA and is presented as present (+) or absent (-).

Neutralizing Mab epitope resides on VP2 of IBDV.

Non-neutralizing Mab epitope resides on VP3 or IBDV.

Recombinant baculovirus vaccines incorporating unmodified large segment A GLS proteins.

Current recombinant baculovirus vaccine incorporating modified chimeric large segment A GLS proteins.

A major difference in the unmodified and chimeric vaccines was the appearance of the classic B69 epitope in the chimeric GLS product. The level of the B69 epitope was arbitrarily set at 9 since no comparisons could be made to the unmodified GLS subunit vaccines. By comparing the status of Mab defined epitopes on the challenge viruses with the unmodified and chimeric GLS subunit vaccines (Table 3), it could be seen that while the chimeric product had expressed the B69 epitope found on the Classic STC and IM challenge viruses, that it also retained all of the homologous GLS epitopes.

Active-cross protection

Table 4 shows the results of a cross-protection trial and serological results obtained prior to challenge.

TABLE 4

Active cross-protection induced 2-weeks post immunization with baculovirus expressed chimeric I-7 IBDV antigens and associated prechallenge serology.

			Nu	Number Protected	þí	Mean VN Titer Log	iter Log	
Group No.	VaccinationA	Challenge ^B	AC-ELISAC	Histo	BBWR ^D	D78	GLS	Mean ELISA
1	None		N/A	NA	NA	54	\$ I	0
Ħ	None	STC	9/0	0/10	0/10	\$ ₹	\$1	0
III	None	MI	9/0	0/5 ⁸	5/5 ⁸	× × ×	\$ FI	0
ΙΛ	None	E/Del	9/0	01/0	0/10	4 4	14	0
>	None	GLS-5	9/0	0/10	0/10	54	5.4	0
VI	1-7	SIC	5/2	10/10	10/10	107.7(1.8)	10.4(1.4)	1235(312)
VII	1-7	MI	5/5	10/10	10/10	10.0(1.4)	10.4(2.1)	1201(791)
VIII	1-7	E/Del	5/2	10/10	10/10	11.4(1.2)	10.6(1.9)	1089(409)
ΙΧ	1-1	GLS-5	5/5	10/10	10/10	11.0(1.5)	12.0(2.0)	1220(339)
×	I-7	None	5/5	NA	NA	9.9(1.4)	9.3(1.4)	1140(473)

Avaccination was given at 8-weeks of age.

*Challenge virus was given by intraocular instillation 3-weeks post immunization or at 11-weeks of age for controls.

Protection was determined by AC-ELISA examination of 1/3 of each group 4-days post-challenge.

Protection was determined histologically and by bursa to body weight ratios at 8-days.

 $^{ ext{ iny F}}$ ive chickens were scored as dead due to IM challenge prior to 8-days post-challenge.

One standard deviation.

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Groups II - V served as challenge controls and as indicated by AC-ELISA, bursa to body weight and histological assessments, all non-vaccinated chickens were fully susceptible to virulent IBDV challenge with all strains used. The IM challenge produced lethal disease in one-third of the control group chicks. In contrast, 8-week old chickens comprising Groups VI - IX were vaccinated once with the GLS chimeric vaccine, and 3-weeks PI all vaccinated chickens were completely protected from challenge by all challenge viruses, including lethal disease produced in controls by the IM strain. Serologically, titers from reciprocal-cross VN tests conducted on prechallenge sera with the D78 and GLS tissue culture viruses were essentially within 2-fold of one another. Mean ELISA titers were relatively low, but were also uniform between the vaccinated groups.

Characterization of vaccines

In initial studies with Baculovirus expressed subunit GLS vaccines, after administration of two doses, the V-IBDV-7 GLS vaccine (Table 3) could only induce active antibody levels capable of providing 79% protection against homologous GLS challenge (Vakharia et al (1993) "J. Gen. Virol.", 74:1201-1206). In a subsequent study, the antigenic mass of the original V-IBDV-7 vaccine was increased approximately 4-fold (calculated at the group specific Mab 8 site) and initiated one dose and two dose vaccination cross-challenge trials with the unmodified GLS subunit vaccine designated as V-IBDV-7-1 (Table 3). In those trials, two doses of the vaccine yielded complete cross-protection against virulent STC, E/DEL and GLS challenge. However, in the one vaccine dose trial, while complete protection was attained against challenge with variant E/DEL and GLS viruses, only 44% protection was achieved against the more distantly related Classic STC virus. Those studies could mean that simply by increasing the antigenic mass and/or doses of the vaccine that better crossprotection could be obtained. However, it was also evident in the absence of homologous vaccination that lower levels of

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antibody, induced by one dose of the GLS V-1BDV-7-1 subunit vaccine, were not sufficiently cross-protective against Classic IBDV challenge. This could mean that in even lower levels of antibody, such as in cases of waning maternal antibody, that cross-protection would likely be even more reduced. Indeed, although not challenged with the STC virus, in some passive maternal antibody studies conducted using another dosage of the V-1BDV-7 vaccine, while homologous GLS protection was afforded, progeny of vaccinated hens were only 57% protected against a more closely related E/DEL challenge.

In a single-dose vaccination cross-challenge trial, the chimeric GLS I-7 vaccine, which incorporated the Classic B69 neutralization epitope, was evaluated. In order to make the current trial comparable to previous trials, the I-7 vaccine was formulated such that by AC-ELISA the relative antigenic mass of the I-7 chimeric subunit vaccine was nearly identical to the unmodified V-IBDV-7-1 vaccine previously used (Table Table 4 shows the results of the cross-challenge after a single dose of the I-7 vaccine was administered. Results were similar to those obtained with the unmodified V-IBDV-7-1 vaccine previously used in that protection against the GLS and E/DEL strains was complete. However, the I-7 vaccine yielded complete protection against pathogenic and lethal challenge by the Classic STC and IM strains respectively. Since the antigenic mass of the GLS and group common epitopes on V-IBDV-7 and I-7 vaccines were carefully equilibrated and equal, it is reasonable to conclude that the comparative increase in efficacy of the I-7 vaccine against challenge with Classic IBDV strains was due solely to the incorporation of the Classic IBDV B69 neutralization epitope in the GLS VP2 protein sequence.

VIRUS-LIKE PARTICLES

As noted above, the recombinant cDNA and immunogens expressed thereby, of this invention may be confined to the VP2 immunogenic region. In other words, it may be sufficient to prepare a cDNA clone encoding epitopic determinants for a

base IBDV, e.g., GLS, as well as a second IBDV epitopic determinant, such as D78. Other epitopic determinants, all in the VP2 epitopic determinant region may be incorporated, cloned and expressed as discussed above.

As reflected in Figure 2, virus-like particles are generated by the expression of DNA encoding the VP2-VP4-VP3 structural protein sequences. These virus-like particle immunogens can be separated from the corresponding VP2 only immunogens, both in terms of monoclonal antibody and by conventional separation measures, such as electrophoresis and chromatography. The difference in reactivity with monoclonal antibody strongly indicates, however, that epitopes present in the VP2-VP4-VP3 structural protein sequence-induced virus-like particles are present that are not present in immunogens expressed by the identical VP2 only region. These epitopes are "both linear and conformational" epitopes. Conformational epitopes differ from linear epitopes and are reflected in the conformation, not only in amino acid sequence of the actual virus. As a result, inoculation of poultry with a recombinant virus-like particle may provide even superior protection against field challenge from IBDV than inoculation with the immunogens of the VP2 region only. This is due to the spontaneous assembly of all the structural elements of the virus.

Applicants have discovered that the expression of the VP2 region as part of the VP2-VP4-VP3 structural protein single segment generates virus-like particles such as those of Figure 2. These particles have been demonstrated to react with antibodies which do not react similarly with the identical recombinant VP2 immunogen. Thus, the virus-like particles may give rise to higher antibody titers, and/or subtly different (broader) protection when a poultry host is inoculated therewith.

The invention herein therefore embraces (1) recombinant VP2 immunogens comprising epitopic determinants of at least two different IBDV strains and (2) virus-like particles of VP2-VP4-VP3 segments wherein the VP2 region again comprises

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epitopic determinants of at least two different IBDV strain, as well as the nucleotide sequences encoding both 1 and 2, and vaccines embracing the same.

RECOMBINANT EPITOPIC DETERMINANT COMBINATIONS:

As reflected in the examples set forth above, genetic epitopic determinants for an IBDV strain can be inserted in the VP2 region of a different, base IBDV genetic sequence, and subsequently used to express an immunogen exhibiting epitopes for both IBDV. Indeed, the examples above demonstrate the combination of at least three different IBDV epitopic determinants. More can be combined. The resulting vaccine includes an active agent, the expressed immunogen, which provides challenge protection against a broad spectrum of IBDV, rather than prior art virus-based vaccines which give protection against a single strain, or a single family of strains.

Figure 7 reflects the amino acid identities for the epitopic determinant region for seven different IBDV. These are not intended to be limiting, but are representative. Desirable recombinant immunogens, both VP2 only and virus-like particle VP2-VP4-VP3 immunogens are made by substituting the genetic epitopic determinants for the varying amino acids at the identified locations in Figure 7 (locations not identified are conserved throughout the IBDV strains). This induces the expression of the inventive immunogens. Clearly, the possible combinations, while large in number, are limited, and may be investigated with routine skill. Representative combinations will tend to reflect combinations of epitopic determinants for dominant IBDV.

A E/Del/GLS recombinant may include changes in the E/Del epitopic determinant region at position 213, Asn-Asp, 253 Gln-His and 169 Thr Ser.

A DS326/D78 recombinant may include the amino acid, and corresponding nucleotide substitutions at 76Ser-Gly, 249 Lys-Gln, 253 Gln-His and 270 Ala-Thr substitutions.

Obviously, a wide variety of combinations are possible

and will occur to those of skill in the art. The epitopic determinant region, roughly including the region from amino acid 5-433 of the VP2 region, thus constitutes a recombinant "cassette" which may be tailored by site-specific mutagenesis to achieve amino acid insertion and/or deletion to provide desired recombinant cDNA clones, polypeptides, virus-like particles and vaccines with improved protection against a wide variety of IBDV.

LETHAL IBDV, MONOCLONAL ANTIBODY AND VACCINE THEREFORE

As noted, typically, IBDV infection creates an immunosuppressive condition, and is reflected in lesions in the bursa of Fabricius. This is typical of IBDV countered in the United States. There exist, however, lethal IBDV, that is, IBDV infections which results in chicken mortality directly as a result of IBDV infection. While vaccines have been developed on the basis of isolation of these IBDV, the resulting vaccines are "hot", that is, they themselves create or induce an immunosuppressive condition, and the inoculated chick must be bolstered with antibodies to other infectious agents. This method of protection is so undesirable as to have been discontinued in most commercial poultry houses in Europe. No adequate safe vaccine against the lethal IBDV is currently available.

The inventors have developed a monoclonal antibody, Mab 21, deposited under Budapest Treaty conditions at the American Type Culture Collection, Deposit Accession No. ATCC HB 11566. This monoclonal antibody is specific and neutralizing for lethal IBDV strains. The specificity is reflected in Table 5, which confirms that unlike other monoclonal antibody, Mab 21 is specific for an epitope exhibited only by IBDV strains having lethal potential.

		TABLE 3	ŀ									
Source	IBDV Strain	Coment	8 <u>7</u>	©)	<u>2</u>	의	13 1	S	糽	1 9	27	되
	Lethal Potential											
	+==		+	+	+	+	+	+	+			
Sherma			+	+	+	+	+	+	+			
USDA	STC		+	+	+	+	+	+	+			
Spafas	2512 (Winterfield)		+	+	+	+	+	+	+			
Edgar	Edgar	(vaccine (hot)	+	+	+	+	+	+	+			
•	Pathogenic Virus											
Sterwin	Bursa Vac	(vaccine hot)	+	+	+	+	+	+	+			•
	Vaccine Virus											
ASL	Univex-BD	(ST 14)	+	+	+	+	+ .	+			,	
Select	Bursal Disease Vaccine	(Luk)	+	+	+	+	+	+				
Select		(STD + VAR)	+	+	+	+	+	+				
Kev Vet	Bio-Burs 1	(078)	+	+	+	+	+	+				
Key Vet	Bio-Bus V	(Luk)	+	+	+	+	+	+				
Key Vet	Kev-Burs	(078)	+	+	+	+	+	+	,			
TO TO	Mary and	(Naster seed)	+	+	+	+	+	+	•	,		
100	NAC	(Basendale	+	+	+	+	+	;				
19:10:40	- 87CF		+	+	+	+	+	;		•		
Sterwin		(1 ab Strain)	+	+	+	+	+	-/+				,
Lukert	EAG ((2512)	4	•	+	+	+	. +	•			
CEVA	Bursa Blend	(2)(2)		. ,				. 4	,		•	
InterVet	920		٠						ı		•	
InterVet	Prime Vac		+	+	+	+	+	+		•	+	•
InterVet	8903		+	+	+	•	+			+		
Solvav	Bursine	(Luk)	+	+	+	+	+	+	•			
Solvay	Bursine 11	(Luk+)	+	+	+	+	+	+				١.
•	Lab Virus											
JKR	E/bel		+	+	+		+			+		
JKR	A/Del		+	+	+		+			+		
KKR	D/De1		+	+	+		+	•		+		
088	GLS		+	+	+	+					+	
DBS	DS326		+	+		+			ı		+	+
*Skeels	24 S	(Serotype II)	+	+	+	+	+	+				
2			+	+		+	+	+		\cdot		
* Field Strains: A	 Field Strains: All classic filed strains tested to date which NOTE: 1. Lulert and STC are Edgar derivatives. 2. Univax is a 	tested to date which were isolated in the U.S. have the 21 marker s. 2. Univax is a Bursa Vac derivative. 3. Bursa Blend is a 2512 Winterfield derivative.	have the arsa Bler	21 ma dis e	rker 2512 v	interfi	eld de	ivativ				

It should be noted that throughout this application, reference is made to a variety of monoclonal antibody which are used to confirm the presence of epitopes of different IBDV in the inventive recombinant chimeric immunogens of the application. These monoclonal antibody have also been deposited under Budapest Treaty conditions and are freely available. They are not, however, necessary for the practice of this invention, and do not constitute an aspect thereof. This should be contrasted with Mab 21.

Like other Mab developed by the inventors herein for IBDV, passive immunization against IBDV lethal strains, particularly designed to achieve immunization in a uniform, standardized level, and to augment any maternally derived levels against lethal IBDV field infection can be obtained by vaccinating one-day old chicks with a vaccine comprising a pharmacologically acceptable carrier such as those described above, in which is present an amount of Mab 21 effective to provide enhanced protection for the inoculated chicks.

The necessary level of protection can be conferred to by a single dose of the vaccine administered in ova or to a day-old chick having a Mab 21 concentration of between 1 microgram and 1 milligram, or repeated vaccinations having a smaller effective dose, but carried out over time. If repeated vaccinations are used, the dosage levels should range between 1 microgram and 1 milligram. The concentration level needed to vaccinate older chickens increases with the weight of the bird and can be determined empirically.

Further investigation of the amino acid sequences of the lethal strains in the epitopic determinant region reflects the highly conserved 279 identity Asn at position 279 of VP2, in non-lethal strains, with a conserved Asp identity at the same position in lethal strains. Accordingly, the lethal strain epitopic determinant recognized by Mab 21, unique to the lethal strains, empirically differs from non-lethal IBDV by the substitution 279 Asp-Asn. According to the methods set forth above, a chimeric, recombinant immunogen conferring effective protection against lethal IBDV, something not

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possible previously with <u>any</u> type of vaccine without inducing an immunosuppressive condition, may be prepared by inserting the genetic epitopic determinant for 279 Asp in a non-lethal base IBDV, such as GLS. This will confer protection against the base IBDV, the lethal IBDV, as well as all other IBDV whose genetic epitopic determinants are inserted. Vaccines prepared from these immunogens, whether VP2 only, or in the form of virus-like particles of VP2-VP-VP3 segments, are used in the same fashion as discussed above.

Claims:

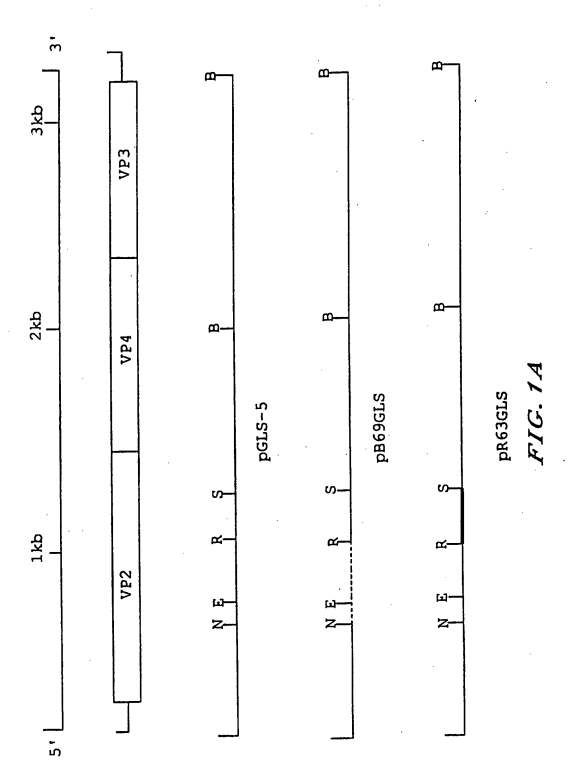
- 1. A chimeric polypeptide immunogen comprising the VP2 amino acid sequence of a first infectious bursal disease virus (IBDV) except for at least one amino acid X, wherein X is an epitopic determinant from a second IBDV strain.
- 2. The immunogen of Claim 1, wherein said VP2 amino acid sequence comprises a plurality of a different epitopic determinant X.
- 3. The immunogen of Claim 2, wherein said plurality of epitopic determinants X are from at least two different IBDV strains.
- 4. The immunogen of Claim 1, wherein said IBDV strains are selected from the group consisting of GLS, E/Del, D78, DS326, RS593, Cu-1, PBG98, 52/70, STC and 002-73.
- 5. The immunogen of Claim 1, wherein said immunogen comprises the amino acid sequence, in order, for IBDV structural proteins VP2-VP4-VP3.
- 6. The immunogen of Claim 5, wherein said immunogen is in the form of a virus-like particle.
- 7. The immunogen of Claim 6, wherein said immunogen exhibits at least one IBDV conformational epitope.
- 8. The immunogen of Claim 1, wherein said amino acid sequence includes an epitopic determinant X of a lethal IBDV strain.
- 9. The immunogen of Claim 8, wherein said epitopic determinant of lethal IBDV strains comprises the amino acid Asp at position 279 of the VP2 sequence.

- 10. A preparation sufficient to provide poultry inoculated therewith resistance to IBDV challenge from at least two different IBDV strains, comprising, as an active agent, an effective amount of the immunogen of any one of Claims 1-9, and a pharmacologically acceptable carrier.
- 11. An avirulent immunogen which confirms on poultry inoculated therewith protection against challenge from IBDV lethal strains, said immunogen comprising the VP2 amino acid sequence of an IBDV, wherein position 279 of said VP2 amino acid is Asp.
- 12. The immunogen of Claim 11, wherein said immunogen comprises, in order, amino acid sequences for VP2-VP4-VP3 IBDV structural proteins.
- 13. The immunogen of Claim 12, in the form of virus-like particles.
- 14. A monoclonal antibody which binds, under AC-ELISA conditions, to IBDV lethal strains, and has the epitope binding characteristics of the monoclonal antibody expressed by the cell line deposited under Accession No. ATCC HB 11566.
- 15. The monoclonal antibody of Claim 14, wherein said monoclonal antibody is obtained, directly or indirectly, from said cell line.
- 16. The monoclonal antibody of Claim 15, wherein said antibody is the antibody expressed by said cell line.
- 17. A preparation for conferring passive immunity in a poultry inoculated therewith against IBDV lethal strain challenge, comprising, as an effective agent, the monoclonal antibody of any one of Claims 14-16 in an effective amount, and
 - a physiologically acceptable carrier.

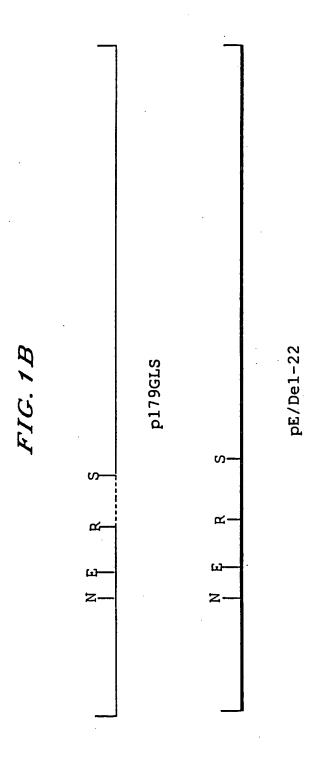
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- 18. A chimeric cDNA which, when operably inserted as heterologous DNA in the DNA of an expression host, encodes the immunogen of any one of Claims 1-9.
- 19. A transfection vehicle for the infection of an expression host, comprising the cDNA of Claim 18 as operably connected in the DNA of baculovirus fowlpox virus, turkey herpes virus or adenovirus.
- 20. An expression vehicle for the expression of the immunogen of Claims 1-9, comprising an expression host selected from the group consisting of SF9 cells, chicken embryo fibroblast cells, chicken embryo kidney cells and vero cells transfected with the vehicle of Claim 19.



SUBSTITUTE SHEET (RULE 26)



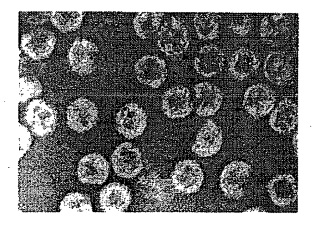


FIG.2A

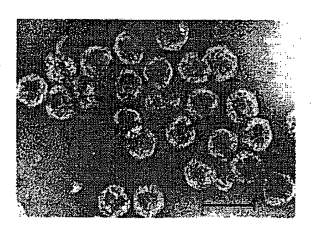


FIG.2B

					4/38						
09	OTGSGLIVFF						L - 120 -	1 1		1 1	1 1
50					1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1				; ; ; ; ; ; ; ; ; ; ; ; ; ; ; ; ; ; ;		
40			#		1						
30					1	100					
50	RSLLM 	XXX	 			06	KFD 				
10	T00 			8				S			X
	GLS DS326 E/Del	D78 Cu-1 PRG9	52/7 crc						1 1 1 1 1 1 1 1 1 1 1		- A
	20 30 40 50 		GLS MTNLQDQTQQ IVPFIRSLLM PTTGPASIPD DTLEKHTLRS ETSTYNLTVG DTGSGLIV DS326H	GLS MTNLQDQTQQ IVPFIRSLLM PTTGPASIPD DTLEKHTLRS ETSTYNLTVG DTGSGLIV DS326	GLS MTNLQDQTQQ IVPFIRSLLM PTTGPASIPD DTLEKHTLRS ETSTYNLTVG DTGSGLIV D78	GLS MTNLQDQTQQ IVPFIRSLLM PTTGPASIPD DTLEKHTLRS ETSTYNLTVG DTGSGLIVFF DS326	GLS MTNLQDQTQQ IVPFIRSLLM PTTGPASIPD DTLEKHTLRS ETSTYNLTVG DTGSGLIVFF DS326 E/Del D78 Cu-1 PBG98 XXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXX	10	GLS MTNLQDQTQQ IVPFIRSLLM PTTGPASIPD DTLEKHTLRS ETSTYNLTVG DTGSGLIVFF DS326	GLS MTNLQDQTQQ IVPFIRSLLM PTTGPASIPD DTLEKHTLRS ETSTYNLTVG DTGSGLIVFF BD326	GLS MTNLQDQTQQ IVPFIRSLLM PTTGPASIPD DTLEKHTLRS ETSTYNLTVG DTGSGLIVFF DS326 E/De1 D78 Cu-1 PBG98 XXXXXXXXXXXXXXXXX OH D70

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	09	PFNLVIPTNE	ATG - 360 -A 361
	20	GLTTGTDNLMA-I	LVAYERV
<u>ن</u>	40	VITRAVAANN	110 NYPGALRPVT
FIG. 3C	30	YLIGFDGSA TT TT TT TT TT -VTT	3 GSLAVTIHGG
	20	HSLVLGAT Q QG QG QG QG QG	90 BGDQMSWSA, A-E A A A A A A
	10	SVGGELVF KTSV	BIVTSKSGG KD D- D-
		GLS S DS326 - E/Del D78 - Cu-1 PBG98 - 52/70 STC - O02-73 -	TATISTITATE SHEET (RULE 26)

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60 	YL - 480
50 ILSERDRLGI	10 LA HAIGEGVDYL
40 DPGAMNYTKL	TLFPPAAP]
FIG. 3D 30	IRRIAVP
20 	80 90
SVVTVAGVSN	DLSSPLKINNNNN
GES GES GES DO S Cu - 1 PBG98 52/70 STC OH	70 DFREYFMEVA

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60 VVDGILASPG	I - 600
ANLEQVPQNP \ ANLEQVPQNP \	OPPSQRGSFI
40 	00
AASGRIRQLT LAAL	TPKALNSKI
20 	80 90
10 LGDEAQAASG	VLREGATL
GLS DS326 E/De1 D78 Cu-1 Cu-1 PBG98 52/70 S7C OO2-73	70 ILRGAHNLDC V V V V V V V V V V V V V V V V V V V

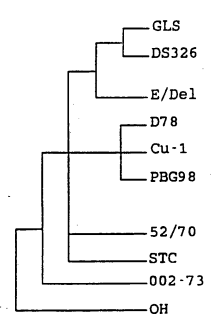
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$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	9/38	80 VAMTGALNAC G	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
യമാത്ര	Cu-1 PBG98 52/70 STC 002-7 OH	70 DVFRPKVPIH	, , , , 1 1 1 1 1 1 1 1
GL DS D7	SUBSTITUTE SHEET (I	* *	iiii

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09	MEAAASVDP N N N N N N N	- 840
50	ETPELESAVR A	AKYGTAGYGV
3 <i>G</i>	HLAMAASEFK E7	110 APQAGSKSQR
FIG. 3G	VLPPNAGRQY HJ	100
50	WDRLPYLNLP Y.	90
10	IKRFPHNPRD W	80 WLEENGIVTD
	GLS DS326 E/Del D78 Cu-1 Cu-1 SBG98 52/70 STC O02-73	THOSALSVEM
	and the state of	Mule 20)

20	RGPSPGQLKY WQNTREIPDP	INHGRGPNQE - 960
FIG. 3H 40	EARGPTPEEA OREKDTRISK KMETMGIYFA TPEWVALNGH R	B0 90 110
	GLS EARC DS326 E/Del D78 Cu-1 PBG98 52/70 STC OO2-73	70 NEDYLDYVHA EH

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		H	1	•	1	1	1	!	1	i	1	
	50	PTORPPGRLG RWIRTVSDED LE	1 1 1 1 1 1 1	1 1 1 1 1 1	1 1 1 1 1 1 1		1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	1 1 1 1 1 1 1	1 1 1 1 1 1 1 1	1 1 1 1 1 1 1 1	! ! ! ! !	
.31	40	PTORPPGRLG	1 1 1 1 1 1 1 1 1	1 1 1 1 1 1 1 1	t 1 1 1 1 1 1 1	 	t 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	1 1 1 1 1 1	1 1 1 1 1 1 1 1 1 1	-818-	S	
FIG. 31	30	PPKPKPRPNA	X	K	LK	LK	LK	K	X	K	K	
	20	EMKHRNPRRA	1 1 1 1 1 1 1 1	1 1 1 1 1 1 1 1 1 1	1 1 1 1 1	i I I I I I I	1 1 1 1 1 1	1 1 1 1 1 1 1 1 1	T-	 	 	
	10	QMKDLLLTAM EMKHRNPRRA PPKPKPRPNA PT	1 1 1 1 1 1 1	1 1 1 1	1 1 1 1 5	1 1 1 1 1 1 1	1 1 1 1 1 1	1 1 1 1 1 1 1 1	1 1 1 1 1 1 1 1 1	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	1 1 1 1 1 1 1 1	
		GLS	DS326	E/Del	D78	Cu-1	PBG98	52/10	STC	002-73	НО	
		SI	BS							HII	F 9	í

FIG. 4



Done on large genome segment A of GLS-IBDV.

DE From cDNA clones pGLS-1 to pGLS-4.

Total number of bases is: 3230. Analysis done on bases 114 to 3152. Done on (absolute) phase(s): 3. Using the Universal genetic code.

14/38 ච්ච චච CCC GCG CCC GGG GAC AGG CCG TCA AGG CCT TGT TCC AGG ATG GAA CTC GCA Ser CGG TGG TTA GTA GAG ATC GGA CAA ACG ATC 160 ATA 50 TTC 100 150 GTT Val ATT Gln CAG CAA 140 CAA ACC TGA GAT Gln Asp TAT CAT CAA CTGLeu ACA AAC (Thr Asn 1 GAG TCA CTA CAA 120 GGG ATG MET CLL

GTA CAC TAC CTA GTG AGT CGG AGT CTC ACA Leu Val Ser Arg Ser Leu Thr GAG AAG His GCC Glu Lys G1yATA GAC ACA Asp Thr GGT GCT Gly Ala ACT Thr ACC CIG Leu Thr Leu CTG Phe Asp Gln MET Leu GGG G1yTIC GAT CAG ATG CIC GTG Val ATT (Ile GAĊ Asn Asp GTG Val 260 420 Leu GGC TCA Asp Thr Tyr Asn Leu Thr Gly Ser GAC TTG ACT. GCA Pro . CCG 470 FIG. 5B 360 AGG Arg Pro CCI GGA CCG GCG TCC ATT Gly Pro Ala Ser Ile ACC TAC AAT TAT Tyr \mathbf{TGC} Cys Lys Gly Phe AAG 410 Val 300 $_{
m Gly}$ TAC Tyr TAC ggg GGA Asn Tyr 460 GGT GCC AGC TAC AAC ACA CTG CAG AGC AAT GGG AAC Thr Leu Gln Ser Asn Gly Asn ACC TCG Thr Ser CCT 350 Ala Ser Tyr Phe TTC400 Len GAG Glu Phe TTT290 180 ACA ACA Ser GIC TCA Val 340 ATT Leu Pro Pro, CCA AGG Arg 230 CTA GGG CTA Gly Leu CIC Leu MET 280 Asn AAC AGG Arg Thr 440 CTG Len 170 330 SUBSTITUTE SHEET (RULE 26)

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GAC Asp Leu GAA Glu TTG Leu Gln AGG GGG GGG Gly Gly 650 GCA ACA Ala Thr GTG AAT Val Val TAT GAT Asp Asp Leu Tyr TAC CTA 590 GAT Val GGG Gly GIC AGC Ser Asn Val 640 Leu AAC CLL Lys MET Leu AAA ATG CTG GTT800 530 Val ACC TAT GAT Tyr Asp ATT GGG Glu Leu Thr Asp ACA GAT Ile Gly 580 Pro ACT 740 ACA Lys GAC TCA Ser GAA CTG AAC GAC AAA Leu Asp Val Thr 790 GTA CTT Asn Asp ACA Thr 680 CCC GAC AGG CCC AGA GTC TAC ACA GGT GGG Pro G1ySer CCC GCT ATA GGG AGT 730 Val ATC I Leu CTG Gly Leu TTA 620 780 510 GGA AGC (Gly Ser I CAA ACA Pro Arg Asn AAC CTC AGC Leu Ser Pro Ala GCC 560 Ile Arg TAC dgg GTT ACT GTC Glv Val Thr Val GAC CCC ATA Gln MET Ser Ala Thr GCA ACA CAA Pro CAG Gln TTC 770 Gly Val Ser ACC Thr ATG TCT 550 AGC GTG Val 009 SUBSTITUTE SHEET (RULE 26)

ACC GGC ACC GAC AAT Thr Gly Thr Asp Asn CAG Gln GTC ATC ACA GGC AAC Gly Asn Ile GGG GAC Gly Asp AGC Ser TCT Ser 920 Pro ACA Thr GGG Asp Gly Gln Glu CAG GAA LysGly AAA GGT 1130 860 1020 CAG Gln His Thr Phe ACC CAT Phe TTTGGA GAG CTC GTG TTT ATA GGC 7 Gly CTG ACG Asn Gly Leu Thr GAG ATA GTG ACG ATT Gly Glu Leu Val 1070 TCC AAA AGT GGT GGT 096 Glu Gly Thr 850 Val Ser Lys Ser ACC AAC Thr Asn GGG Tyr Leu 1010 FIG. 5D TAC AAT AGC CTA GCA Ser Leu Ala 1060 CCA Pro ' GGG AAC Asn Gly Ile 950 Thr GCA Ile ACC ACC ATT GLL Val 1000 ATA GTG Ile Val Ser Ala GIG GTG 999 CTG GGC GCC Leu Gly Ala Gly AGC gcc Leu Val 890 1050 Val CLI AGT GTG CIC Ser Leu 940 AGC Ser Arg Ala Phe Asn Leu Glu Ala CTG GAG GCA TTC AAT AGA GCT 1100 830 TCG Ser GAT GCC ATC ACA GTA Ile Thr Leu Val 880 Lys Pro Trp Thr TGG CITACT CCA ICC ATC AAA Ile MET Ser ATC Ser Asp Ala 820 Leu CAC Val CTT 980 870 SUBSTITUTE SHEET (RULE 26)

Leu Trp Gly Ala MET Asn CCA AAT CCT GAA Pro Asn Pro Glu GTG GCA ACA GGA Val Ala Thr Gly CGG Val Thr ACA GGA GGC Ile Lys Glu Val ATC AAG TAC GAA AGA GTG 1400 GAG GTG Pro AAA GAC Lys Asp GAG CTG ATC Glu Leu Ile Tyr Glu Arg CCA 1180 GGC gly ATG MET Glu Leu TTT GAC Gly Arg Phe Asp 1340 TTC Phe TIC CTG AGT GAG AGG GAC CGC CTT Tyr Thr Lys Leu Ile Leu Ser Glu Arg Asp Arg Leu Glu Tyr Phe 1390 GAG TAC TTT GGC Phe Gly ACA CTA GTA GCC Thr Leu Val Ala GGG GTG AGC AAC TTC Ser Asn Phe GGC CGA 1280 1330 GGA GCA GIY Ala CCG ACA AGG GAG TAC ACC GAC TTT CGT Pro Thr Arg Glu Tyr Thr Asp Phe Arg GAA TAC Leu Ala Lys Asn Leu Val Thr Glu Tyr 1380 Gly Val 1270 ATT GCA Ile Ala Val CTG GTT ACA CCC GIC GTC GCT (Val Ala GGG GCC CTC CGT CCC Gly Ala Leu Arg Pro 1210 CCC CTG AAG Lys TTG ATA Arg Glu Tyr 1260 Thr Pro Leu ACG GCA AAG AAC 1420 1150 TAC ACA AAA GTTVal Val Ser 1360 AGC Ser Ser CTA 1250 1410 SUBSTITUTE SHEET (RULE 26)

GCC TCA GGC Ala Ser Gly Asn Leu CCA CCT GCC GCT CCC Pro Pro Ala Ala Pro GAG GCA CAG Gly Glu Ala Gln 1570 Pro GIC GCG Val Ala TCA CCC 1620 GGC GCC GIY Ala Ser 1510 GCT GCT CGA GCC GCG TCA GGA AAA GCA AGG GCT Ala Arg Ala Ala Ser Gly Lys Ala Arg Ala GGT GAT Gly Asp GCC GCC GAC AAG GGG TAC GAG GTA Ala Ala Asp Lys Gly Tyr Glu Val Gly Ile Leu Ala 1670 1560 CTT Glu GAG 1720 TTC GGG GAA GGT GTA GAC TAC CTG CTG Gly Glu Gly Val Asp Tyr Leu Leu Leu Phe GGG ATT Leu Arg 1610 1500 TTA TTG Thr Leu Ala Ala Asp Lys Gly 1660 GTC GAC (Val Asp (GTG Ser Thr Val TCC ACA 1550 1710 TGC Cys 1600 GTG GTC Pro Val CIC GAC CCC GTA Leu Asp 1490 Pro Val Val 1650 1540 CCG AAC Asn ACT CTC CAG AAT Gln Asn 1700 CAC GTGHis Val 1480 1750 Ala GGA ACT G Pro GCA ATT Ala Ile Ala CCC Ile Arg Gln Leu GCA GCT CGC ATA AGG CAG CTG 1640 1530 GTG GGT Gly Gln Val 1690 CAG CAT GCT GCT TCA Ala Ala Ser CGC Leu Ala His ATA AGG AGG Arg Arg Leu Arg 1580 1470 GCC TIC Phe Ala Ala 1630 CIG Arg CTA Leu 1520 1680

CCC AAA CTG TCC AAA GAC CCC ATA CCT CCT ATT Leu Ser Lys Asp Pro Ile Pro Pro Ile GCA CTA AAC Ala Leu Asn Ser GCT TCT 1840 Pro GTT GTC CCA Val Val Pro TyrTAT CCA Pro Pro 2000 1890 GGA ' Arg CGA CCI 2050 TTT Phe ATG ACA CCC AAA MET Thr Pro Lys Glu Asp Leu Gln TyrCAA 1940 GTC TAT 2100 1830 GGG AGA GAC TAC ACC Gly Arg Asp Tyr Thr Trp Asp Asp Ser Ile MET Leu Ser Lys Asp Pro CIC GTG Val GGA CAC AGA GTC Gly His Arg Val 1990 Asp GAT GAG GAC 1380 2040 ATG Tyr GCT TAC GAA GGC GTG CGA Glu Gly Val Arg GTC ATC ACG ACA GTG GAA GAC GCC Val Ile Thr Thr Val Glu Asp Ala 2090 1820 1980 Ala ACT CTC TCC Thr Leu Ser 1870 AGT GGA AAC CTA GCC ATA Ser Gly Asn Leu Ala Ile GAG ACT Glu Thr TGG GAC GAC AGC ATT ATG 2030 1920 2080 1810 GTC ATT C GAT GGG GTA CTT CCA CTG Gly Val Leu Pro Leu TCC TTC ATA CGA Ile Arg 1970 1860 2020 Phe Ala Ser Phe 1910 2070 1800 1960 GIC AAC Gly Val GGA Asn Val Ser Lys MET GTG AAA ATG 1850 2010 Asp Asp Pro . GGA Gly CAA AGA GAT CCT Gln Arg 1900 AGC CCA Pro Asp GAT GTG TIC Phe Val 2060 1790

GCA Phe CTC CCC TFC CTC AAC Leu Pro Tyr Leu Asn Lys GAG AAG GGC CTC 7 Gly Leu 1 GAA GCC ATG GCC MET Ala GAG ATT TGG GCA ACG 2270 2160 MET Trp Ala 2320 CTC GCC I gcc Ala CAC CGG CTT His Arg Leu Cys Gly TGT GGC 2210 AGG AAC ACC GGG CCC AAC Asn Thr Gly Pro Asn 2260 GAA CTC GAG AGC GCC GTC Glu Leu Glu Ser Ala Val GAC TGG GAC AGG His Asn Pro Arg Asp Trp Asp Arg TAC CAC Tyr His ACG GGA GCC CTC AAC GCT Thr Gly Ala Leu Asn Ala 2420 2150 2310 AAG CTC GCC ACC GCA Lys Leu Ala Thr Ala 2200 GGA CGC CAG Gly Arg Gln 2360 FIG. 5H 2250 2410 CIC CCA CGC GAT GTA Gly Pro Gly Ala Phe Asp Val 2300 2190 GCA Pro Asn Ala 2350 TTC CCT CAC AAT CCC AAT ACC CCT Thr Pro TTT AAA ATA AGC TTT AGA AGC ACC Lvs Ile Ser Phe Arg Ser.Thr GCC ATG 2240 Ala MET 2400 2130 GCT GGT CCC GGA GCA 2290 Glu GAG Leu Pro Phe Pro CTT CCA Ile Ser Phe Arg GTG His Val 2180 2340 CAT AAG Lys GAG TTC CGT CCA TAC Pro Tyr Glu Phe Pro Ile Lys Arg 2390 CCC ATC 2120 2280 AAA Ala Leu Leu CLILys TTG GIC Val 2330 2220

GAG ACC GGC CCC ACA GAA GAG AAT GGG ATT GTG ACT GAC ATG GCC AAC TTC GCA CTC AGC GAC CCG AAC Glu Glu Asn Gly Ile Val Thr Asp MET Ala Asn Phe Ala Leu Ser Asp Pro Asn CIG TCT Leu 2650 Gly Pro MET Glu His Arg GCC CAT CGG ATG CGA AAC TTT CTT GCA AAC GCA CCA CAA GCA GGT AGC AAG Ala His Arg MET Arg Asn Phe Leu Ala Asn Ala Pro Gln Ala Gly Ser Lys \mathbf{I} GG 2540 ATG (ATG MET 2590 CGG AAG AAG Arg TIC Lys Lys AAT GGG Asn Gly Phe 2750 2480 2640 GAG GCC (Glu Ala A AGT GTG Val 2530 2690 TCA Leu CTC Ser Ser Val Asp Pro Leu Phe Gln Ser Ala Leu Ser GTA GCA Val Ala CGG ATC Arg Ile TIC CAA ICI GCA CIC TAC GGA GTG Gly Tyr Gly Val 2740 2470 2630 2520 TGG GAA AAA GAC ACA Trp Glu Lys Asp Thr 2680 Glu GAA GGG ACA GCA GGC 2570 2730 2460 CCA (Pro (Gly Thr Ala 2620 ACA Thr GCA GCC AGT GTA GAC CCA CTG 2510 2670 GCA AGG 2560 CAG Gln Phe TAC Tyr TAC TTT 2720 2610 AAA . GCA Lys 2500 AGG GCC A GAA GAA (Glu Glu ATC 2660 2550 CAA AGG (Gln Arg 7 GGC Ala Ala 2440 ATG MET Pro 2600 2490

MET GCI GAG Pro Gly Gln Ala Glu Ser Glu CGG TTG GCA TCA GAA 2920 CAT GAG CCC AAC Pro Asn CAG GCA CAG AAC ACA CGA GAA ATA CCG GAC Asn His Glu Ile Pro Asp 3080 2970 MET ATG GAA ATC AAC Arg Leu Ala 2860 GGA Glu Ile ATG AAA GAT CTG CTC TTG ACT GCG MET Ivs Asp Leu Leu Thr Ala CCA CCA AAG CCC AAG CCA AGA Pro Lys Pro Arg 3020 2910 CCA 3070 2800 AAG AGC GCTTyrTyr Leu Asp Tyr Val His Ala Glu Lys Ser Gly Ala GTC TAT Gln Asn Thr Arg 2960 2850 999 FIG. 57 Pro Asn Gln Glu Gln MET I.ys Asp Leu Leu Glu Val Ala Lys Val 3010 Pro Lys GCT TTC ATA GAC GAA GTT GCC AAA GCA GAG CTA AGG GCA GCT ACG TCG ATC TAC Ala Ala Thr Ser Ile Tyr 2900 3060 2950 Pro TAC TGG Tyr Trp 2840 TAT CTA GAC TAC GTG CAT 3000 GCT (Ala] 2890 CGT GGC CCA AAC CAA GAA CAG Ile Asp AAG CAT CGC AAT CCC AGG CGG Arg Asn Pro Arg Arg Gln Leu Lys 3050 2780 CAG CTA AAG 2940 2830 Leu Arg Pro Gln Ala Phe 2990 2880 CCC GGC (Pro Gly (3040 2770 Ile GAG GAC Glu Asp CAA ATC CCA CCC CAA 2930 2820 AGC ($_{
m G1y}$ His Gln Ser 2980 Glu (CCA Pro Arg AAC Pro Asn GAA 2870 3030 2760

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rc rcr gar al Ser Asp	3190	CA GGC GTG		·	PC/Gene
AGG ACT GARG Thr V	3180	ACC CGC 0			B(
c rgg Arc g Trp Ile	0-	c gac acc	3230	G GAT CCG	
cTG GGC CG Leu Gly Ar	317	SAG TCT CC	3220	ccc aaa tr	
GGT CGG (3160		10	CAA CAT	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
3A CCC CCT rg Pro Pro	3150	AG TGA GGC lu	32	CG GCC TTA	06
cca acg cag a Pro Thr Gln A		asg gac ctt g 3lu Asp Leu G	3200	SAC ACC AAT T	2 Aug - 1990
	des cressed and and act of the Arg Len Gly Arg Trp Ile Arg Thr Val Ser	CTG GGC CGC TGG ATC AGG ACT GTC TCT Leu Gly Arg Trp Ile Arg Thr Val Ser 3170 3180 31	ACG CAG AGA CCC CCT GGT CGG CTG GGC CGC TGG ATC AGG ACT GTC TCT GAT Thr Gln Arg Pro Pro Gly Arg Leu Gly Arg Trp Ile Arg Thr Val Ser Asp 3150 3160 3170 3180 3190 GAC CTT GAG TGC TCC TGG GAG TCT CCC GAC ACC CGC GCA GGC GTG Asp Leu Glu	ACG CAG AGA CCC CCT GGT CGG Thr Gln Arg Pro Pro Gly Arg 3150 GAC CTT GAG TGA GGC TCC TGG ASP Leu Glu 3200 3200 3210	ACG CAG AGA CCC CCT GGT CGG CTG GGC CGC TGG ATC AGG ACT GTC TCT GAT Thr Gln Arg Pro Pro Gly Arg Leu Gly Arg Trp Ile Arg Thr Val Ser Asp 3150 3160 3170 3180 3190 Asp Leu Glu 3200 3220 3220 3230 ACC AAT TCG GCC TTA CAA CAT CCC AAA TTG GAT CCG

************************* A NUCLEIC ACID SEQUENCE

Done on DNA sequence EDEL22

E/DEL virus, vero cells adapted

Analysis done on the complete sequence

genetic code

ACG GAG ATC AGA GTA TGG TTA TGA CAT Total number of bases is: 3180.

Analysis done on the complete second on (absolute) phase(s): 1.

Done on (absolute) phase(s): 1.

Using the Universal genetic code

10 20

GAA TTC CTC CTT CTA CAA CGC TAT TAT

ATA Ile GTT CCG 1 CAC CAG ATT His Gln Ile 150 CAA ACC Gln Thr 140 GAT Gln Asp CAA ACA AAC CTG Thr Asn Leu ATG 120 gag GCA ATC 110

Leu GAC GAC ACC Asp Asp CCG Pro Ile TCC Ser Pro Ala Gly ACA ACC Thr Thr CCA ATG CTG Leu Leu CTTAGC Ser CGG

AAC

CTA

TAT

ggc

GGT

CCT

CTC Leu

ACA

Ser

Ser

AGG

Thr

GTA

ACA

CIC

380

Val

Asn

Leu GGŤ CTC GIG Val ATG Ser Gln MET AGT grĠ Val GIG Val GGC TCA ACT Gly Ser 260 Leu Asp TAC AAT TTG Tyr Asn Leu GAT Arg $\mathtt{T}\mathtt{T}\mathtt{C}$ AGG 200 TGC Lys AAG TTC 360 FIG. 6B 250 TAC TAC TyrGGA $_{
m G1y}$ Asn AAC Pro AAC Asn ACC TCG 300 TTC Gly TAC TyrGGGSer AGC GAG AGT Ser 240 Val CTG CAG AGC GIC TCA Ser Gln Ser 290 Leu Pro AGG Arg 180 Leu CTA Leu GGG CTA Gly Leu CTCAsn AAC CAG TAC Tyr Ser TCA Lys His AAG CAC Ğly **GGG** CAC His CCC Ala 330 AC'L Thr GAG

GCC GAT AAT Ala Asp Asn Asp Asn TAC CTA GTT AGC Val Ser GTC Val CTT CGG Asp Leu Gly GGG AAC C AAA ATG Lys MET 480 640 GCA GAT Asp GAT 530 CCA ACA (Pro Ile Thr TAT ACT Thr ATC AAC GAC AAA ATT 069 580 TCA CTT GAC Leu Asp ATA Ile CTG Ile Asn Asp Lys Glu Leu Ser ACC Thr GAA ACA Thr630 520 GTC TAC CCC GGG Gly Pro GGA AGC CTG AGT Ser Leu Ser 680 ATA (Ile (Leu TTA 570 Asn Ser CCC AGA AAC AGC CCC GCT Pro Ala Arg Pro Arg 620 CCC CTCGly Ala Leu 510 670 AGG ACC GTC Thr Val ATA CAA Gln ACA 560 GAC CCC A 450 TTC GAC GCA Ala Asp GTA TCT Ser AGT ACC Thr Ser 500 GAA GGG (Glu Gly TTG ATG AGC GTG GGTGlyVal Ser 099 550 GAC CCC Asp CLL Len ATA AAC Ile Asn Arg $_{
m TGT}$ Cys AAC GGG Asn Gly AGG G1y009 490 GTG ACA Thr Val GTA 650

AAA Lys GAT TCA GGC TTT Gly Phe TTC 860 CTG ACG Len CTG GAG CTC (ACA Thr ATC Ile ATC TAC CTT Ile Tyr Leu Asn GGG AAT 960 850 ACC GGA Asn Asn AAT GGG GTA ACA Gly Val Thr GTT GGG (Val Gly (AAC Pro CCA 900 GCA ACC Thr ATT 950 GTG CTC AGC (Leu Ser GCC GGT 840 CAA ACA (Glu Thr (GGC $_{
m G1y}$ Phe Asn Leu GCT GTG Arg Ala Val 890 CIG Leu TTC AAT Ser AGT 780 AGA CTT GTA TAC GAT GCC ATC ACA Asp Ala Ile Thr Leu Val ATC ACC A CCA CAG Gln 720 AGC Ser ATG TCA Asp Ala CAA Leu CTT GTA Val 930 Ser GTC AAT Asn AGC Asp ACT Thr Ala Asn 760 GGG Gly ATC Ile Thr 920

GGA CCA Pro G1yCAG GCA Gln Ala GTG Val 1080 GGT Gly Pro CCA His CAT GAA AGA Glu Arg 1130 Asp GAC Leu CTG ATC Ile G1yGGT1020 1180 Glu Phe GAG TTT TAC TyrThr Ser Asp ACG GAT 1070 Phe Val ĠCC Ala TTCCGA GTG 1230 Gly Arg AGT 1120 gga Asn AAC GIG Val GCA Ser Leu Ala Ser Lys TCC AAA 1010 TAC CTASer Leu GTG AGC 1170 AGC CTA 1060 GAA Thr Val Thr Glu ACA ACC 1220 FIG.6EACA ggg Gly GGG GIC Gly GAG ATA GTG Glu Ile Val 1110 Pro Val GLT מממ GCT AGT Ser Ala Val 1160 GIC CTG Leu Val GCC CTC CGT Ala Leu Arg TCG GCA Ser Ala 1210 Asn AAC Thr ACG Len CIG1100 AAG TrpAla Lys GTC GTT Ile TGG Val Val ATA 990 1150 Gly GGA Ser ATC ATG TCA Ile 1040 Glu Leu Ser CTA MET CCA Pro \mathbf{ICI} TCC Ser 1090 GAA GGA Tyr ATC ACA Ile Thr CAG Gln TAT 980 GCA ACA Ala Thr CCT Pro AAC Asn Glu GAA 1140 1030 1190 AAT Asn GGC GGG CCA Gly

CTG GGC GAT Lys1350 AAA GAG ATA GAG GTG ATC AAG Glu Val Phe Pro TTG TTC CCA Lys Asp 1400 Leu MET ATA CTG AGT GAG AGG GAC CAC CTT GGC Ile Leu Ser Glu Arq Asp His Leu Gly ATG 1290 Leu Phe Leu TIC CTA GAG TAC TTC Glu Tyr Phe 1340 TyrGGC Ser Thr TyrPhe Gly GAC TAC TCT ACA 1500 1390 Asp Glu Glu Arg Asp 1280 Val GTG GTC Gly Ala GGA GCA TTT CGT Phe Arg Pro Val Val 1330 Glu Gly GAA GGT FIG. 6F 1270 CCC GCA GAC Thr Asp Pro Leu Lys Ile Ala Ile Leu Ser 1380 GGG GTA Ile Ala Val GCI TAC CCC CTG AAG Glu Tyr 1320 GCA TTG GAG GCC ATA AGG AGG ATA Leu 1370 CAT CCA ACA AGG Tyr Thr Lys Pro Thr Arg Ala Ile Arg Arg TAC ACG AAA Leu Asn Ser CTC AAC TCT 1310 Pro Leu CCT CTA 1470 Trp $_{
m LGG}$ Ala MET Asn GCC ATG AAC Arg GCT GCC GAC Als Asp ATC CGG rhr Val 1410 1300 Iel 1460 GCC ACC

AAA Lys GAG GTA Glu Val GCT 1620 CCC Glu GAG CTT Leu AGG Arg 1670 AAA GCA Lys Ala AGA ACA Thr ATT TAC TyrLeu Arg 1560 CTA GGG G1yATG MET GGG Gly 1610 GCC GIG GGA AAG GAC Asp G1yLys Val Asp Ala 1660 \mathbf{TGC} GAC Cys GTC Val GAC Asp GCG TCA Ala Ser 1550 Glu GCC Ala GAA GTA CTC GAC Asn Leu Asp Ala Pro Val 1600 gcc Ala \mathtt{GTG} CCC Ala gcc Val 1760 FIG. 6G Thr AAC Leu AAT ACA Gln Asn CGA CIC Thr Ala Arg 1650 CAC His ACG Thr CAG Thr ACT GCT Ala Pro Len ATT GCA ACC CTG מממ 1590 Gly GTC Gln GGT GGA Gly CAG GIG Gln Val 1640 CAG CGC GIG AGG Arg Arg Val TCA Ser 1530 Leu ATA TTCPhe GCTAla CTT CCT 1580 CGC Arg Leu ATA TTC CTA GCT 1740 1630 999 Leu Gly CTA CAG Gln GGC G1yAAT Asn 1520 ACG GCG CCC Pro Thr Ser GCA 1680 1570 1730 GCC Ser TCA Ala GIC Val GAG GCC

Phe Gln 1890 TAT CAA Val 2050 GTC CIC GTGLeu TAC ACC Val Tyr Thr 1940 AAG GAC GAT Lys Asp Gly His Arg GAA GAC Glu Asp GGA CAC AGA 1830 1990 ATG GAC Asp ' 1880 GCT TAC A TCC GGG AGA Gly Arg Ser Arg CGA 2040 1930 Leu CIG Glu Gly Val CIC ICC GGC GTG Thr Leu Ser 1820 ACT Glu Thr TGG GAC GAC AGC ATT ATG Ile MET 1980 1870 GAG 1 CTA GCC ATA CGA ACT Leu Ala GTC ATT GAA 2030 Ser Ile Arg GGG GTA CTT CCA CTG Val Ile 1920 1810 Gly Asn GGA AAC Trp Asp Asp 1970 TCC TTC Ala Ser Phe GCI 1860 2020 AGT Ser Phe LLL AAC Asn Gly TCT CAA AGA GGA GAT GTC AAA ATG Ile Asp Asp Val 1800 Lys MET 1960 GGA Gly Ala Pro Asp Gln Arg GCT CCA GAT 1850 GCA CTG AAC AGC GTC CCA ATA GAT Ser 2010 ATT GTG Val Ile Ser Leu Asn Tyr TAT CCA Pro Pro Pro CCT CCT 1840 2000 Pro

Leu

Tyr

Gln

Arg

G1y

Pro Asn Ala

Leu Pro CTT CCA

CGC CAG

GGA

AAT GCA

CCCC

TAC Tyr

CCA

Leu Asn Leu

TAC CTC AAC CTT Tyr Leu Asn Leu

2270

2300

CAC

CCC Pro TGG Leu 2160 Leu \mathbf{TGT} CCC AAC Pro Asn Arg CGG AAG CTC GCC ACC GCA CAC Lys Leu Ala Thr Ala His Asp Arg GAC AGG Asn Ala 2100 CTC AAC GCT 9999 Gly 2150 TrpLeu TGG ACC Val Asn Thr 2200 Asp GAC GGA GCC Gly Ala GTA AAC 2090 Pro Arg CCA CGC 2140 GAT Asp ACG (Thr FIG. 6IGCC ATG Ala MET CCT CAC AAT Pro His Asn ACC Ser Thr TTC GGA GCA TTC Gly Ala Phe 2190 2080 AGC Val GTGTTC AGA Phe Arg 2130 CAT (His TIC Phe Gly Pro GCT GGT CCC Ser Lys Arg GAG AAA ATA AGC AAA CGT CCC ATC 2070 Pro Glu Lys Ile 2120 GGC CTC AAG TTG Gly Leu Lys Leu TTC ATC GIC CTC AAG TTG Pro Lys Val 2170 Ala Thr Phe CCC AAA GCA ACG ATT SUBSITUTE SHEET CGA

Arg CGG AGC GGC GCC 2430 TTCSer 2590 Ala gaa GCA Val GCA CTC AGA \mathtt{GTG} Leu Arg 2480 Glu Ala CAA Gln GAG Ser AGT GIC Val 2370 2530 Pro CCA GCC (Ala TIC GIG CICPhe Val Ala Leu 2420 GCA GCC AAT Ala Asn GCA G1yAGC Ser GGA 2580 2470 Leu Ala Asn TAC Tyr Ser CTT GCA AAC CTC GAG Glu Leu Glu \mathbf{ICI} 2360 gga Gly MET ATG CAA Val Asp Pro Leu Phe Gln 2520 2410 TIC GAC Asp ACA GCA GAA Thr Ala 2570 FIG. 67 TTTPhe CCT GTG GCT Thr Pro CCA CTG 2460 Val Ala 2350 999 Gly ACC CGA AAT Arg Asn 2510 TAC Glu GTG GAC Tyr ATT GAG 2400 2560 AAG Lys GGG Lys ATG MET AAA 2450 ggg Ala Ala Asn GAG AAT GCA GCC AAT CAT CGG His Arg Glu Asn GAG TTT Glu Phe 2340 2500 Arg TCG CAA AGG 2390 Ala Gln TCA AAC GCC Asn Ala Ser CTG GAA Leu Glu 2550 2440 Ser GCA Ala GCA Ala 2330 Glu Trp Pro AAG Lys GCT Ala GAA GAC CCG TGG 2490 2380 2540 AGC . Asp Ser ATG MET ATG ATG MET MET

ATC TCA AAG AAG CTC AAT GGG Arg CGA GAA Lys Leu Asn AAG AGC CGG 2750 Ser Thr Arg Ser 2800 Ile ACA Lys GGG G1yGCA Trp Val Ala 2690 CAG AAC Gln Asn C C C C GAG Glu Arg CCA GAA TGG GTA ATC TAC 2850 GCA Ala Ile CAG AGG GAA AAA GAC ACA Gln Arg Glu Lys Asp Thr 2630 Ser Pro Glu Val His TAC TGG Tyr Trp GTG CAT 2790 2680 GCT ACG Thr TAC Tyr Ala ACA Thr GGG CCA AGC CCC GGC CAG CTA AAG 2730 Gln Leu Lys GCA CTA GAC GCA Leu Asp 2780 ATC CTA AAG Ile Leu Lys ATG GGC ATC TAC TTT MET Gly Ile Tyr Phe Leu Lys 2670 TyrPro Gly Glu Ala CCG GAC CCA AAC GAG GAC TAT GAA GCA 2720 Glu Asp 2610 CAA Glu CCA GAG Ser Gln 2660 Pro (Gly Pro Pro Asn 2820 GAA GAA Glu Glu ACC GGC CCC ACA ThrPro Thr 2600 GAG Glu CGA Arg Pro Asp TCA Ser 2650 2810 CAC (His / Gly GCAAAla MET

2970 GCA GAA ATC Arg ATC AGG ACC ACC ACT Thr 3020 Leu AAG Pro Lys GTC TAT Tyr 2910 3070 GAC Leu Leu CTG CTC Val 2960 AAG CGC Lys Pro Lys TCT CCC Gly Arg 3120 AAA 3010 GGC CCA Asp GAT GAA GTT GCC Glu Val Ala 2900 Pro GGG CAG ATG AAA Gln MET Lys CCA CTG Leu 3060 2950 CGG Gly Arg FIG. 6T CCC ICG GCT 3110 GGT GAC Asp 3000 CGG Arg Asn Pro Arg Arg 2890 GGG Glo ATA CCC AGG GAA CCC CCT Pro Pro 3050 TTC Phe Pro Asn Gln CAA 2940 Arg GAG AGA CGC AAC GCT Ala CCT AAC 2990 CAG Gln Leu CII CAA 2880 3040 His 1 GAC ACA GGC G1yCAT Thr Glu Asp Pro CCC 2930 Pro GAG Arg CCA Pro CGTAAG MET Lys 3090 CCA 2980 GAT Asp ATG GCT GAG Glu 2870 Ser GAG Glu Pro Asn \mathbf{ICI} His CCC AAT GCA CAT 3030 2920 3080 Asn] GIC AAC ATG MET CAG

TTG GAT CCG FIG.6MGCA GGC GTG

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NS	2,	263	ren	=	=	<u> </u>	=	=		Phe	ng.	
AMINO ACID CHANGES IN VP2 VARIOUS IBDV STRAINS	IN VP	258	Gly	E	#	2		=	F	2	Asn	
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(SOS)	AMINO ACID RESIDUE NUMBER IN VP2	253	His	듄	2	His	2	2	등	=	2	
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S N		242	lo/	=	2	2	E	2	<u></u>	ΙOΛ	2	
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		76	Ser	r	=	G	Ser	2	2	2	R	
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	330	Ser	2	=	Arg	Lys	Arg	Ser	=	
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VIRUSES	•	CLS	SD326	E/0EL	078	Cu-1	PBC98	52/70	STC	002-73

INTERNATIONAL SEARCH REPORT

International application No. PCT/US95/03772

A. CLASSIFICATION OF SUBJECT MATTER						
IPC(6) :Please See Extra Sheet. US CL :Please See Extra Sheet.						
According to International Patent Classification (IPC) or to both national classification and IPC						
B. FIELDS SEARCHED						
Minimum documentation searched (classification system follow	ed by classification symbols)					
U.S. : 424/159.1, 185.1, 186.1, 204.1; 435/320.1, 252.3	; 530/350, 388.3, 397, 402, 403; 935/10	, 12				
Documentation searched other than minimum documentation to	he extent that such documents are included	in the fields searched				
Electronic data base consulted during the international search (name of data base and, where practicable	, search terms used)				
APS, MEDLINE, EMBASE, CA, BIOSIS, CABA SEARCH TERMS: IBDV, INFECTIOUS BURSAL DISEA ACID, VACCIN?, DNA	SE VIRUS, VP2, VP4, VP3, ANTIBO	D?, ASP, ASPARTIC				
C. DOCUMENTS CONSIDERED TO BE RELEVANT						
Category* Citation of document, with indication, where	appropriate, of the relevant passages	Relevant to claim No.				
X JOURNAL OF GENERAL VIROL	JOURNAL OF GENERAL VIROLOGY, Volume 74, issued					
1993, V.N. Vakharia et al., "Infe						
Y Structural Proteins Expressed in		1				
Confer Protection in Chickens", p	ages 1201-1206, see entire	9				
document.	•					
X ARCHIVES OF VIROLOGY, Volur	ne 120. issued 1991. C.D.	1-4				
- Bayliss et al., "A Recombinant Fo						
Y the VP2 Antigen of Infectiou		8, 10, 18-20				
Protection Against Mortality Car	used by the Virus", pages					
193-205, see entire document.	·					
·	·					
1						
X Further documents are listed in the continuation of Box	C. See patent family annex.					
Special categories of cited documents:	"T" later document published after the inte date and not in conflict with the applica	ernational filing date or priority				
"A" document defining the general state of the art which is not considered to be of particular relevance	principle or theory underlying the inv					
"E" carlier document published on or after the international filing date	"X" document of particular relevance; the considered novel or cannot be considered.					
*L° document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other		Ť				
special reason (as specified)	onsidered to involve an inventive	step when the document is				
 "O" . document referring to an oral disclosure, use, exhibition or other means 	combined with one or more other such being obvious to a person skilled in the					
P document published prior to the international filing date but later than the priority date claimed						
Date of the actual completion of the international search	Date of mailing of the international sea	irch report				
08 JULY 1995	19 JUL 1995					
Name and mailing address of the ISA/US	Authorized officer	1				
Commissioner of Patents and Trademarks Box PCT	ANTHONY C. CAPUTA					
Washington, D.C. 20231 Faccimile No. (703) 305-3230	Telephone No. (703) 308-0196	7 7				

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INTERNATIONAL SEARCH REPORT

International application No. PCT/US95/03772

			Relevant to claim No		
Category*	Citation of document, with indication, where appropriate, of the relevant	nere appropriate, of the relevant passages			
Y	JOURNAL OF GENERAL VIROLOGY, Volume 70, iss 1989, K.J. Fahey et al., "A Conformational Immunogen of Infectious Bursal Disease Virus that Induces Virus-Net Antibodies That Passively Protect Chickens", pages 1473 see entire document.	14-17			
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INTERNATIONAL SEARCH REPORT

International application No. PCT/US95/03772

A. CLASSIFICATION OF SUBJECT MATTER: IPC (6):

A61K 35/76, 39/12, 39/395; C07K 14/005, 16/08; C12N 1/21, 5/10, 15/33

A. CLASSIFICATION OF SUBJECT MATTER: US CL :

424/159.1, 185.1, 186.1, 204.1; 435/320.1, 252.3; 530/350, 388.3, 397, 402, 403; 935/10, 12

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